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MONOVALENT LIGAND OF THE FcalphaRI RECEPTOR AS AN ANTI-INFLAMMATORY AGENT.

The invention relates to the use of a monovalent ligand of the $Fc\alpha RI$ IgA receptor as an anti-inflammatory agent.

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Immunoglobulin A (IgA) is the most heterogeneous Ig isotype in humans, existing in multiple molecular forms such as monomeric, polymeric and secretory IgA; it comprises two subclasses IgA1 and IgA2.

In serum, IgA exists mainly in monomeric form, with a minor percentage of polymeric IgA (pIgA).

secretions In mucosal (saliva, tears, colostrum, gastrointestinal fluids, nasal bronchial secretion, and urine), IgA is produced as dimers, joined by a polypeptide termed J-chain. Dimeric IgA binds to the membrane-associated polymeric Ig receptor (pIgR), and the resulting complex is transported from the baso-lateral to the apical/luminal side of mucosal epithelium. this transport the bound IgA is released by proteolytic cleavage from the pIgR; however a portion of the pIgR, the secretory component, remains associated with dimeric IgA, forming altogether secretory IgA (SIgA).

SIGA plays a major role in the innate immune system preventing microorganisms and foreign proteins from penetrating the mucosal surfaces. It also neutralizes toxins and infectious organisms.

role the of secretory Whereas is established in mucosal immunology, the function of serum IgA antibodies is mostly unknown. Although IgA is the second most abundant Ig isotype in serum, it is not usually involved in humoral immune responses and does not complement. Monomeric serum activate IqA has inflammatory activity and is capable of down-regulating functions such as IgG-induced phagocytosis, bactericidal activity, oxidative burst, and cytokine release. contrast, polymeric IgA and IgA-containing immune

complexes (IC) can efficiently trigger immune effector functions on blood leukocytes through IgA Fc receptors.

Receptors for the Fc region of immunoglobulins (FcRs) play a major part in the link between humoral and cellular responses. FcRs for all five human antibody classes have been described.

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The human IgA Fc receptors (Fc α R) family comprises several members (for review cf. MONTEIRO and VAN DE WINKEL, Annu. Rev. Immunol. 21: 177-204, 2003), but only Fc α RI (or CD89), a receptor specific for the IgA Fc region, has been identified on blood myeloid cells (MONTEIRO and al., J. Exp. Med. 171: 597-613, 1990; MALISZEWSKI and al., J. Exp. Med. 172: 1665-1672, 1990). Fc α RI is expressed on monocyte/macrophages, dendritic cells, Kupffer cells, neutrophils and eosiniphils and binds both IgAl and IgA2 (CONLEY and DELACROIX, Ann. Int. Med. 106: 892-899, 1987; KERR, Annu. Rev. Immunol. 12: 63-84, 1994) with low affinity (Ka \approx 10⁶ M⁻¹) (MONTEIRO and VAN DE WINKEL, 2003, aforementioned).

Fc α RI is a member of the Ig gene superfamily. It comprises two extracellular Ig-like domains (EC1 and EC2), a transmembrane region and a cytoplasmic tail devoid of recognized signaling motifs. Crystal structures of human FcαRI reveal that the two Ig-like domains are 25 oriented at right angles to each other and that two $Fc\alpha RI$ molecules are required for the binding of molecule (HERR and al., J. Mol. Biol. 327: 645-657, 2003). The IgA binding site is located in the membranedistal EC1 domain. Anti-FcaRI mouse and human monoclonal antibodies (mAb) have been generated (MONTEIRO and al., 30 Immunol. 148: 1764-1770, 1992; SHEN et al., J. 4117-4122, 1989; PCT WO 91/05805; 143, Immunol. PCT WO 02/064634), and it has been shown that monoclonal antibodies that bind in the EC1 domain of FcoRI block IgA 35 binding, whereas those that bind in EC2 do not.

Due to the moderately fast on- and off-rates of the FcaRI:IgA binding reaction, monomeric IgA binding is transient, whereas polymeric IgA and IgA immune complexes bind with a respectively growing avidity due to a decrease in the off-rate (HERR and al., 2003, aforementioned; WINES, J. Immunol. 162: 2146-2153, 1999).

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The involvement of $Fc\alpha RI$ in the ability of IgAresponses such trigger immune as phagocytosis, antibody-dependent cytotoxicity (ADCC), cell-mediated superoxide generation, cytokine production, presentation and inflammatory mediator release, has been reported (for review, see MONTEIRO and VAN DE WINKEL, 2003, aforementioned). It has been proposed to use anti-FcαRI antibodies, such as My 43 (PCT WO 91/05805), or the monoclonal antibodies disclosed in human PCT WO 02/064634, to activate these FcaRI-mediated immune responses.

It has also been proposed to use anti-FcR antibodies, including anti-Fc α RI antibodies, as vectors for selectively targeting active principles, such as cytotoxic compounds, to cells expressing Fc receptors (PCT WO 99/41285).

US patent 6 018 031 describes bifunctional antibodies containing the binding region of an anti-Fc α R antibody and the binding region of an antibody directed against a target cell. These bifunctional antibodies can bind on one hand said target cell, and on the other hand effector cells expressing Fc α R. Their binding to Fc α R triggers the Fc α R-mediated activity of the effector cell, resulting in the destruction of the target cell bound to the same bifunctional antibody molecule.

Signaling through FC α RI is dependent on association of FC α RI with the FCR γ chain subunit, forming the trimer FC α RI $\alpha/\gamma\gamma$. The FCR γ chain contains an immunoreceptor tyrosine-based activation motif (ITAM) in its cytoplasmic tail (PFEFFERKORN and YEAMAN, J. Immunol.

153: 3228-3236, 1994; LAUNAY and al., J. Biol. Chem. 274: 7216-7225, 1999) that allows the recruitment of crucial signalling effectors (KINET, Annu. Rev. Immunol. 17: 931-972, 1999). Fc α RI can be expressed with or without physical association with FcR γ subunit. The γ -less Fc α RI internalises and recycles IgA to the cell surface, whereas $FcR\gamma$ -associated $Fc\alpha I$ directs complexed IgA to lysosomes (LAUNAY and al., 1999, aforementioned; SHEN and al., Blood 97: 205-213, 2001). No cellular function of non aggregated FcaRI, other than IgA recycling, has so far been identified. Receptor aggregation is required for FcaRI-mediated activation of target cell functions such as cytokine release and antigen presentation (SHEN and al., 2001, aforementioned; PATRY and al., Immunol. 86: 1-5, 1995; GEISSMANN and al., J. Immunol. 166: 346-352, 2001).

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While involvement of FcαRI in IgA-mediated inflammation is well recognized, the molecular basis that underlies the IgA anti-inflammatory capacity has not been elucidated until now. Although it has been reported (WILTON, Clin. Exp. Immunol. 34, 423-8 1978; VAN EPPS and WILLIAMS, J Exp Med 144, 1227-42 1976) that IgA inhibitory functions require the Fcα region, the part played by IgA Fc receptors remains unknown.

A consensus model of negative signaling in the 25 immune system involves receptors with an immunoreceptor inhibitory motif tyrosine-based (MITI) in their cytoplasmic domain. These inhibitory receptors act by coaggregating with activatory receptors: cross-talk between the two receptors generates a negative signal (RAVETCH 30 and LANIER, Science, 290, 84-89, 2000). An example of the ITIM class of inhibitory receptors is the Fc\u03c4 receptor FcyRIIB. However no ITIM receptor for the Fca region is known.

The Inventors now found that unexpectedly, monomeric occupancy of FcαRI by a monovalent Fab fragment

of an antibody directed against the EC2 domain of Fc α RI strongly inhibited IgG-induced phagocytosis and IgE-mediated exocytosis, in vitro, and that, surprisingly, these effects were mediated by the ITAM motif of the Fc α RI-associated FcR γ subunit.

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Further, the Inventors have shown in an asthma model, that in vivo targeting of Fc α RI by said monovalent Fab fragment abolished antigen-induced bronchial hyperreactivity and the accompanying airway inflammation particularly leukocyte infiltration into the lung tissue. They have also shown in a model of interstitial renal fibrosis and obstructive nephropathy, that in vivo targeting of Fc α RI by said monovalent Fab fragment considerably decreased the pathological inflammatory reactions.

An object of the present invention is the use of a monovalent antibody fragment directed against the EC2 domain of the Fc α RI receptor, as anti-inflammatory active principle in the preparation of a medicament for treating an inflammatory disease.

The anti-inflammatory properties of said monovalent antibody fragment result from a down-regulation of the pathological inflammatory reactions involving $Fc\alpha RI$ -expressing myeloid cells.

Examples of inflammatory diseases that can be treated according to the invention include allergic diseases in particular asthma, as well as inflammatory diseases involving interactions between immunoglobulins and FcR, such as nephritis, rheumatoid arthritis and auto-immune diseases (lupus, diabetes, etc). They also include non-immune inflammatory diseases such as those induced by unilateral ureteral obstruction resulting in kidney inflammation, drug induced toxicity of the kidney, gut inflammatory disorders such as Crohn's disease.

A monovalent antibody fragment is an immunoglobulin fragment that has only one antigen-binding

site, in contrast with a whole immunoglobulin molecule, that comprises at least two antigen-binding sites. Examples of monovalent fragments are Fab fragments that consist of the light chain and the first half of the heavy chain, or scFv fragments that consist of the variable portions of the heavy and light chains of an antibody, connected to one another via a flexible linker (CLACKSON et al., Nature, 352, 624-628, 1991), thus forming a single-chain protein.

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Methods allowing to obtain monovalent antibody fragments that can be used in the practice of the invention are well known in themselves.

way of example, Fab fragments can be by the conventional techniques of enzyme obtained, digestion, from an antibody directed against the EC2 domain of the $Fc\alpha RI$ receptor. Said antibody can be a murine monoclonal antibody obtained by the conventional hybridoma technology. Advantageously, it can also be a chimeric antibody, a humanized antibody, or a completely human antibody. Chimeric antibodies can be obtained from said monoclonal antibodies by replacing the constantregion domains by human domains; humanized antibodies can be obtained by incorporating the CDRs of said monoclonal antibodies into the framework regions (FRs) of a human antibody, using techniques, known in themselves, of CDR grafting. Completely human monoclonal antibodies can be the way as conventional obtained in same monoclonal antibodies, except that the mice immunized are transgenic mice with a human immunoglobulin repertoire, as disclosed for instance in PCT WO 02/064634.

Monovalent antibody fragments, in particular scFv fragments, can be directly obtained by expressing, in an appropriate host cell, a recombinant DNA comprising the DNA sequences encoding the variable regions of a monoclonal, humanized or human antibody directed against the EC2 domain of the Fc α RI receptor, associated with an

appropriate linker. They can also be generated from an antibody phage display library, panned with the EC2 domain of the $Fc\alpha RI$ receptor. Humanized scFv fragments can also be obtained by the method described by ARNDT et al, (Int J Cancer 107, 822-829, 2003).

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The specificity towards the EC2 domain of the FcαRI receptor of the above antibodies and monovalent fragments can be checked by testing their effect on the binding of IgA to the $Fc\alpha RI$ receptor; the antibodies or fragments that do not block said binding are in most of cases directed against the EC2 domain. However, some nonblocking antibodies such as the monoclonal antibody A3, have been reported to recognize an epitope between EC1 and EC2 domains (MORTON et al., J Exp Med, 189, 1715-22, 1999). Accordingly, the above test will advantageously be completed or replaced by an assay of the binding of said antibodies or monovalent fragments to a recombinant protein comprising the EC2 domain and devoid of the EC1 domain of the $Fc\alpha RI$ receptor, such as the chimeric receptor composed of Fc α RI EC2 and bovine Fc γ 2R EC1 MORTON et al. (1999, cited above). described by Alternatively, the monovalent fragments of anti-Fc α RI antibodies that do not block the binding of IgA to the FcaRI receptor can directly be tested in vitro for their anti-inflammatory properties, for instance their ability to inhibit IgG-mediated phagocytosis in human blood monocytes, or to inhibit the IgE-mediated degranulation response of a mast-cell line expressing FcαRI, described in the examples below.

For the practice of the invention, the monovalent antibody fragments can be administered, systemically or locally, in various ways.

By way of example they can be administered by the parenteral route, including for instance intramuscular, intradermal, intravenous, intraperitoneal, subcutaneous, or local injections.

Local administration in the respiratory tract can also be used, provided that the monovalent antibody fragments of the invention are in a form suitable for delivery to mucosal surfaces of the airways. For example, they may be suspended in a liquid formulation for delivery to a patient in an aerosol form or by means of a nebuliser device similar to those currently employed in the treatment of asthma.

The monovalent antibody fragments can be optionally mixed with suitable carriers and/or excipient(s) known to those of ordinary skill in the art.

The present invention will be understood more clearly from the further description which follows, which refers to non-limiting examples of preparation and of use of monovalent antibody fragments of anti-Fc α RI antibodies in accordance with the invention.

EXAMPLE 1: FCCRI TARGETING INHIBITS IGG-MEDIATED PHAGOCYTOSIS IN HUMAN BLOOD MONOCYTES IN VITRO

The role of $Fc\alpha RI$ in the modulation of IgG-20 mediated phagocytic activity of blood monocytes was examined.

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Human peripheral blood mononuclear cells were Ficoll-Hypaque isolated by density gradient centrifugation from healthy volunteers. Enriched (70 to 80%) monocyte populations were obtained by adherence to described and plastic in MONTEIRO al. as (1990, aforementioned).

Fab fragments of an anti-FcαRI mAb (IgG1κ, clone A77, MONTEIRO and al., J. Immunol. 148: 1764-1770, 1992) and of an irrelevant control monoclonal antibody (IgG1κ, clone 320) (PASTORELLI and al., J. Biol. Chem. 276: 20407-20412, 2001) were generated by pepsin digestion for 8 h at 37°C followed by reduction with 0.01 M cysteine and alkylation with 0.15 M iodoacetamine at pH 7.5. Complete digestion and purity were controlled by SDS-PAGE.

blood mononuclear cells Adherent were preincubated with 10 μ g/ml Fab A77 (c), irrelevant Fab 320 or buffer for 30 min at 37°C. After washing, cells incubated at 37°C for 30 min with Texas-redconjugated E. coli (50 bacterial/cell) (Molecular Probes, Eugène, Oregan), opsonized or not with polyclonal rabbit anti-E. coli IgG antibodies (Molecular Probes) according to the manufacturer's instructions. After washing, slides examined with а confocal mounted and were microscope (LSM 510 Carl Zeiss, Jena, Germany). Overlaid 10 transmission and fluorescence images (mid sections) are panels (a-d) are representative of six shown. The number independent experiments. The (±SD) of mean ingested bacteria per monocyte in six experiments with different healthy donors is shown in Figure 1. It was 15 determined by counting at least three fields in each experiment. The number above the bar corresponds to the mean percentage of inhibition by Fab, calculated as follows: $100 - 100 \times (n \text{ of IgG-opsonized bacteria in the})$ presence of Fab A77 - n of non opsonized bacteria) / (nof IgG opsonized bacteria -n of non opsonized bacteria) in which n indicates the mean number of internalised bacteria.

The results are shown in Figure 1.

Legend of Figure 1:

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Non opsonized bacteria (Non ops)

= buffer

IgG opsonized bacteria (IgG-ops)

 \square = buffer

= anti-Fc α RI Fab A77

= irrelevant Fab 320

* P < 0.02, Student's unpaired t test

The results show that IgG opsonization enhanced $E.\ coli$ phagocytosis by monocytes. Preincubation with anti-Fc α RI Fab A77 fragment inhibited IgG-mediated

phagocytosis by more 80% compared to the irrelevant Fab 320 fragment.

EXAMPLE 2: CHARACTERIZATION OF FCORI INHIBITORY FUNCTION

The inhibitory function of FC α RI was further studied by testing the degranulation response of the rat mast-cell line RBL-2H3 that constitutively expresses the high-affinity receptor IgE (Fc ϵ RI), transfected with wild-type human Fc α RI.

1) Material and methods:

10 Cell transfection:

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Transfection of RBL-2H3 cells was performed as described by LAUNAY and al. (1999, aforementioned): the wild-type human FcαRI construct was cloned into pSRαNEO vector containing a resistance gene to neomycin between restriction sites the 15 XbaI-BamHI and sequence controlled by DNA sequencing. RBL-2H3 cells, maintained as described by ROA and al. (J. Immunol. 159: 2815-2823, with 1997), transfected $15 \mu g$ of DNA were electroporation at 250 V and 1500 μ Fa using an Easyjet⁺ apparatus (Eurogenetec, Seraing, Belgium). 20

resistant for 1 mg/ml G418 Clones were selected for $Fc\alpha RI$ expression by flow cytometry. Cells preincubated with 100 µg human polyclonal IgG (PharMingen, San Diego, California) to block FcyRs before incubation with phycoerythrin-labeled anti-FcαRI (IgG1κ, A59-PE) (MONTEIRO and al., 1992, aforementioned) isotype-matched irrelevant Ab with (Becton an or Dickinson, Bedford, Massachussets). After washing, cells were analysed using a FACScalibur flow cytometer and Dickinson). CellQuest software (Becton One clone expressing human $Fc\alpha RI$ (clone 15.4) was selected for the following experimentations.

Degranulation response

Exocytosis of granular mediators contained in 35 cells was determined by measuring the release of $\beta-$

hexosaminisase as described in (ROA and al., 1997, aforementioned), by $Fc\alpha RI$ transfected cells, or by non transfected cells used as a control, upon sensitization with different test reagents.

Dickinson) at 5×10^4 cells/well. Cells were sensitized with different test reagents as hereafter indicated for each reagent. Cells were washed in prewarmed Tyrode buffer (135 mM NaCl, 5 mM KCl, 5.6 mM glucose, 10 mM HEPES, ph 7.3, 1.8 mM CaCl₂, 1 mM MgCl₂, and 5% BSA), and degranulation was triggered with 0.1 μ g/ml DNP-HAS (Sigma). Net β-hexosaminidase release was calculated as a percentage of total content after subtracting spontaneous release.

2) Inhibition of IgE-mediated exocytosis by anti-FcαRI Fab fragments

Human Fc α RI transfectants (clone 15.4) and non transfected (NT) RBL cells were sensitized with IgE anti-DNP (1:200) or IgE anti-DNP plus 10 μ g/ml irrelevant Fab 320 control or anti-Fc α RI Fab A77 for 1 h at 37°C. Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are shown in Figure 2a.

Legend of Figure 2a:

Wt#15.4 = human Fc α RI transfectants (clone 15.4)

NT = non transfected RBL cells

 $\square = IgE$

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= IgE + A77 anti-FcαRI Fab

= IqE + irrelevant Fab 320

* P < 0.02, Student's unpaired t test

Data are means ±SD of five independent experiments. The number above the bar corresponds to the mean percentage inhibition of degranulation.

The results show that antigen stimulation of IgE-sensitized transfectants (clone 15.4) induced a

strong degranulation response. Preincubation with anti-FcαRI Fab A77 markedly inhibited FcαRI-initiated degranulation (74%), as compared to an irrelevant Fab 320. Similar results were obtained with two others transfectants (not shown) but not with non transfected cells (NT). The inhibitory effect of A77 Fab was even stronger when preincubated for longer periods of time (2 to 12 hours) (not shown).

Of note, anti-Fc α RI Fab failed to modify IgE binding (not shown). Anti-Fc α RI Fab purified by gel filtration had a similar inhibitory action, ruling out a role of aggregates in the observed effects (not shown).

3) Dose response study of anti-FcaRI Fab-mediated inhibition

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE in the presence of different concentrations of anti-Fc α RI Fab A77 of irrelevant Fab 320 for 1 h at 37°C. Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are show in Figure 2b.

Legend of Figure 2b:

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 \blacksquare = IgE + anti-FcαRI Fab A77

= IgE + irrelevant Fab 320

* P < 0.02, Student's unpaired t test.

Data are means $\pm \text{SD}$ of four independent experiments.

The results show that inhibition by anti-Fc αRI Fab was concentration-dependent, and was maximal between 1 and 10 $\mu g/ml$.

4) Influence of epitope targeted by anti-FcαRI Fab on inhibition

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE in the presence of 10 μ g/ml Fab fragment from different anti-Fc α RI mAbs: A3 (recognizing

a binding site between EC1 and EC2; A59, A62, A77, recognizing a binding site within EC2) or irrelevant Fab $320 \text{ for } 1 \text{ h at } 37^{\circ}\text{C}$.

Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are shown in Figure 2c.

Legend of Figure 2c:

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= anti-Fc α RI Fab (A3, A59, A62, A77)

= irrelevant Fab 320

* P < 0.01, Student's unpaired t test

Data are means $\pm \text{SD}$ of three independent experiments.

Three of the four anti-Fc α RI Fab tested inhibited Fc α RI-induced degranulation by >50%. The fourth anti-Fc α RI Fab (A3) failed to inhibit degranulation, even though, like its three counterparts, it bound readily to Fc α RI-transfected cells (not shown).

5) Influence of ligand valence on inhibition

For this purpose, F(ab')₂ were generated from the anti-FcαRI mAb (A77) or from the irrelevant antibody 320, by pepsin digestion for 8 h at 37°C with an enzyme to substrate ratio (w/w) of 1/50 in 0.1 M acetate buffer, pH.4.4 as described in SILVAIN and al. (J. Immunol. 155: 1606-1618, 1995). Complete digestion and purity were controlled by SDS-PAGE.

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE, or IgE plus 10 μ g/ml Fab or F(ab')₂ fragments from A77, or IgE plus irrelevant Fab or F(ab')₂ fragments from 320, for 1 h at 37°C.

Cells were washed, degranulation was triggered with DNP-HSA, and $\beta\text{-hexosaminidase}$ release was determined.

The results are shown in Figure 2d.

Legend of Figure 2d:

 $\square = IgE$

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 \blacksquare = IgE + A77 Fab or F(ab')₂

 \square = IgE + 320 Fab or F(ab')₂

* P < 0.01, Student's unpaired t test

Data are means $\pm \text{SD}$ of four independent experiments.

The results show that monovalent anti-Fc α RI Fab had a stronger inhibitory effect that the divalent F(ab')₂ fragments.

6) Influence of FcaRI aggregation on cell degranulation

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE, or 10 μ g/ml Fab or F(ab')₂ fragments from A77, or irrelevant Fab or F(ab')₂ fragments from 320, for 1 h at 37°C.

Cells were then stimulated with $F(ab')_2$ fragments of rabbit anti-mouse IgG (RAM at 40 μ g/ml) (LAUNAY and al., J. Leukoc. Biol. 63: 636-642, 1998).

Cells were washed and $\beta\text{--hexosaminidase}$ release 20 was determined.

The results are shown in Figure 2e.

Legend of Figure 2e:

 $\square = IgE$

= A77 Fab or F(ab')₂

* P < 0.02, Student's unpaired t test

Data are means ±SD of four independent experiments.

The results show that highly multivalent aggregation of Fc α RI, after crosslinking of anti-Fc α RI F(ab')₂ with rabbit anti-mouse Ig (RAM) F(ab')₂, resulted in degranulation. Less extensive multivalent aggregation with anti-Fc α RI Fab plus RAM F(ab')₂ resulted in weaker degranulation. No degranulation was observed with anti-Fc α RI Fab, F(ab')₂ or with RAM F(ab')₂ alone (not shown).

EXAMPLE 3: SERUM IGA INDUCES FCORI INHIBITORY FUNCTION

The effect of the physiological ligand IgA was tested on Fc α RI RBL-2H3 transfectants (clone 15.4), by testing the degranulation response, as described in Example 2 above.

1) Influence of proteolytic treatment on Fc α RI inhibitory response to IgA

As IgA exert biological activity at inflammatory sites, which contain numerous mediators including proteases, the effect of trypsin treatment of cells on IgA-mediated inhibitory function was examined, given that $Fc\alpha RI$ is resistant to trypsin (MONTEIRO and al., 1990, aforementioned).

Human Fc α RI transfectants were pretreated or not with 1 mg/ml trypsin-TCPK (Sigma) in DMEM for 30 min at 37°C and then sensitized overnight with IgE alone, or with IgE plus 0.2 mg/ml serum IgG, or purified serum IgA (batches n°39328 and 02828, ICN Biomedicals Inc, Aurora, Ohio).

Cells were washed, degranulation was triggered with DNP-HSA, and $\beta\text{-hexosaminidase}$ release was determined.

The results are shown in Figure 3a.

Legend of Figure 3a:

 $\square = IqE$

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= IgE + serum IgA

= IqE + serum IqG

* P < 0.01, Student's unpaired t test

Data are means $\pm SD$ of six independent 30 experiments.

Numbers above the bars indicate the mean percentage of inhibition.

The results show that incubation with serum IgA, but not IgG, significantly inhibited IgE-dependent degranulation (43%). The inhibitory effect of serum IgA,

but not that of IgG, was significantly enhanced ($\sim 50\%$ enhancement) in trypsin-treated cells, while the IgE-mediated degranulation response was not affected. A similar enhancement was observed with purified myeloma IgA (not shown).

2) Influence of Ig concentration on $Fc\alpha RI$ inhibitory function

Human FcαRI transfectants were pretreated with 1 mg/ml trypsin-TCPK (Sigma) in DMEM for 30 min at 37°C and sensitized overnight with IgE alone or with IgE plus various concentrations of two batches of purified serum IgA (batches n°39328 and n°02828, ICN Biomedicals Inc, Aurora, Ohio), secretory IgA (SIgA, batch n°42K3780, Sigma Aldrich, St-Louis, Missouri) or human IgG.

Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are shown in Figure 3b.

Legend of Figure 3b:

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 \Diamond = IgE

= IgE + serum IgA (batch n°39328)

 \triangle = IgE + serum IgA (batch n°02828)

 \bullet = IgE + SIgA

 \Box = IqE + IqG

* P < 0.01, Student's unpaired t test

Data are means $\pm \mathrm{SD}$ of five independent experiments.

The results show that the two different batches of commercial serum IgA inhibited degranulation in a dose-dependent manner, maximal inhibition (66%) being obtained at 0.5 mg/ml. Colostral SIgA also inhibited cell activation, albeit to a somewhat lesser extent.

3) Modulation of FcaRI inhibitory response by IgA1 and IgA2

As $Fc\alpha RI$ binds both IgA1 and IgA2, the inhibitory capacity of the two subclasses was compared relative to that of SIgA which contains variable amounts of both IgA1 and IgA2 depending on the type of secretory mucosa.

Human Fc α RI transfectants were pretreated with 1 mg/ml trypsin-TCPK (Sigma) in DMEM for 30 min at 37°C and sensitized overnight with IgE plus 0.2 mg/ml serum IgG, purified myeloma IgAl and IgA2 or SIgA. Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are shown in Figure 3c.

Legend of Figure 3c:

 \square = IgG

 \blacksquare = IgA1, IgA2 or SIgA

* P < 0.02, Student's unpaired t test

Data are means ±SD of four independent

20 experiments.

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The results show that all tested preparations produced significant inhibition (30-40%) relative to human IgG (<5%). The Fc α RI inhibitory response can be induced by both IgAl and IgA2.

25 4) Comparison of polymeric and monomeric serum IgA inhibition

As Fc α RI binds polymeric IgA more efficiently than monomeric IgA, the inhibitory potential of the various molecular forms of IgA (separated by HPLC), without secondary crosslinking was examined.

Human Fc α RI transfectants were pretreated with 1 mg/ml trypsin-TCPK (Sigma) in DMEM for 30 min at 37°C and sensitized overnight with IgE plus 0.1 mg/ml serum IgG (IgG), total serum IgA (IgA), polymeric serum IgA

(pIgA), dimeric serum IgA (dIgA) or monomeric serum IgA (mIgA). Serum IgA was size-fractionated by HPLC.

Cells were washed, degranulation was triggered with DNP-HSA, and β -hexosaminidase release was determined.

The results are shown in Figure 3d. Legend of Figure 3d:

 \square = IgG

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■ = IgA, pIgA, dIgA or mIgA

inset = size-fractionated serum IgA by HPLC \star P < 0.02, Student's unpaired t test

Data are means $\pm \text{SD}$ of three independent experiments.

The results show that polymeric serum IgA is more inhibitory than monomeric serum IgA. Inhibitory potency increased with the size of the IgA species: polymeric IgA were more efficient (60%) than both dimeric IgA (38%) and monomeric IgA (20%). Similar data were obtained with a different batch of serum IgA separated by HPLC (not shown).

The difference between A77 mAb and IgA may be explained by the binding site and the ligand avidity. While anti-Fc α RI mAb A77 biding site is localized in EC2, IgA interacts with EC1 domain (MORTON and al., J. Exp. Med. 189: 1715-1722, 1999) and polymeric IgA bind more avidly to Fc α RI than monomeric IgA (HERR and al., 2003, aforementioned; WINES and al., 1999, aforementioned). No β -hexosaminidase release was observed when the different IgA preparations were incubated alone with transfected RBL-2H3 cells, and IgE-mediated degranulation was not inhibited in non transfected cells (NT) (not shown).

EXAMPLE 4: THE FC α RI INHIBITORY SIGNAL IS MEDIATED BY THE ITAM MOTIF OF THE FCR γ CHAIN

To explore the structural requirements for the inhibitory signal, a series of $Fc\alpha RI$ mutants and chimeric constructs was used:

– Fc α RI_{R209L} wherein the charged arginine at position 209, within the Fc α RI transmembrane domain, is replaced by a leucine (R209L); this mutation abolishes the association of Fc α RI with the FcR γ chain (LAUNAY and al., 1999, aforementioned; MORTON and al., J. Biol. Chem. 270: 29781-29787, 1995).

- The R209L/ $\gamma_{chimera}$ construct results from the fusion of the extracellular and R209L transmembrane domains of Fc α RI_{R209L} to the intracytoplasmic tail of the human FcRy chain.

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The R209L/ γ_{chimera} was generated as follows. The extracellular and transmembrane domains of the R209L amplified by PCR using primers were mutant GGGCTCGAGATGGACCCCAAACAGACCACC (SEQ ID NO: 1) and $R_{\gamma-\alpha}$: CTTTCGCACTTGGATCTTCAGATTTTCAACCAGTATGGCCAA (SEQ NO: 15 2), as well as the intracellular domain of human $Fc\alpha R$ γ chain using primers $F_{\alpha-\gamma}$: TTGGCCATACTGGTTGAAAATCTGAAGATCCAA GTGCGAAAG (SEQ ID NO: 3) and R_{γ} : GGG**GGATCC**TTACTGTGGTGGTTTC (SEQ ID NO: 4). PCR products were TCATG fused by overlapping extension PCR.

The structures of the wild type $Fc\alpha RI-\gamma_2$ receptor, of the $Fc\alpha RI_{R209L}$ receptor, and of the $R209L/\gamma_{chimera}$ receptor are schematically represented in Figure 4a.

All constructs were cloned into pSRαNeo vector and transfected in RBL-2H3 cells, as described in Example 2.

Cells transfected with wild-type human Fc α RI (clone 15.5), Fc α RI- γ_2 (clone 5.26) or R209L/ $\gamma_{chimera}$ (clone 9.4) construct were selected.

The results of determination of Fc α RI expression by flow cytometry are shown in Figure 4b. These results show that all RBL-2H3 transfectants expressed significant levels of Fc α RI at the cell surface.

degranulation response tested The was as described in Example 2 2).

The results are shown in Figure 4c.

Legend of Figure 4c:

 $\square = IgE$

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= IgE + irrelevant Fab 320

 \blacksquare = IqE + anti-Fc α RI Fab A77

* P < 0.02, Student's unpaired t test

Numbers above the bars indicate the percentage of inhibition as compared to an irrelevant control Fab. 10

show that all transfectants results The sensitised with IgE alone exhibited over 50% Fc&RImediated degranulation. Anti-Fc α RI Fab A77 treatment was non inhibitory in RBL-2H3 transfected with the R209L mutant (clone 5.26) indicating that the intracellular tail of FcaRI did not contain the motif responsible for the inhibitory signaling. In contrast, the binding of anti-Fc α RI Fab A77 to the Fc α RI_{R209L}/ γ chimeric receptor in transfected cells (clone 9.4) restored the inhibitory 20 effect on degranulation to an extent similar to that observed in cells transfected with the wild type receptor (clone 15.5) (91% and 72%, respectively). Similar results were obtained with at least three additional clones for each type of transfectants (not shown).

Aggregation of this $Fc\alpha RI_{R209L}/\gamma$ chimeric 25 receptor induced degranulation, demonstrating that, like wild-type FcαRI, it was able to mediate both activation and inhibition (not shown).

As the FcRy chain does not bear any known inhibitory motif, the FcRy ITAM usually known as 30 activatory motif was investigated to know whether it could also mediate the inhibitory effect. The human FcRy chain contains two carboxy-terminal tyrosines (Y268 and Y278 within the $Fc\alpha RI_{R209L}/\gamma$ chimeric receptor) being part of the ITAM motif known to play a role in cellular 35 activation (17,24). Point mutations (Y268F, Y278F and double Y268/278F) were introduced in ITAM motif of the $Fc\alpha RI_{R209L}/\gamma_{chimera}$.

Stable transfectants (simple or double) established in RBL-2H3 cells transfected with the R209L/ $\gamma_{chimera}$ containing Y268F and/or Y278F mutations within ITAM motif were no longer able to mediate the inhibitory and the activatory response (not shown).

EXAMPLE 5: THE FCORI INHIBITORY SIGNAL INDUCES TYROSINE PHOSPHORYLATION AND AFFECTS Ca²⁺ INFLUX

10 A) Tyrosine phosphorylation assay

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Since ITAM-mediated signalling involves the activation of tyrosine kinases, monomeric targeting of the Fc α RI/ γ complex was investigated to know whether it involved tyrosine phosphorylation.

Indicated RBL transfectants (Fc α RI- γ 2, Fc α RI_{R209L}/ γ _{chimera} wild type, Fc α RI_{R209L}/ γ _{chimera} Y268F/Y278F) were stimulated for 15 min with 10 μ g/ml anti-Fc α RI Fab A77, irrelevant Fab 320, 40 μ g/ml RAM F(ab')₂ or a combination of anti-Fc α RI A77 F(ab')₂ plus RAM F(ab')₂.

After stimulation and two washes in ice-cold PBS, cells were solubilized in lysis buffer (50 mM HEPES pH 7.4, 1% Triton X-100, 0.1% SDS, 50 mM NaF, 50 mM NaCl, 1 mM Na $_3$ VO $_4$, 30 mM Na $_4$ P $_2$ O $_7$, 50 U/ml aprotinin, 10 μ g/ml leupeptin) and post-nuclear supernatants were prepared.

Lysates were resolved by SDS - 10% PAGE and proteins were transferred onto PVDF membrane. After blocking in 4% BSA, membranes were incubated with 4G10 anti-PY Ab (Upstate Biotechnology, Lake Placid, NY) for 1 h at room temperature and with goat anti-mouse Ig coupled to HRP

(Southern Biotechnology Associates, Birmingham, AL). Membranes were then striped and re-probed with anti-rat phospholipid scramblase (PLSCR) mAb (PASTORELLI and al., 2001, aforementioned) to evaluate equal loading. Filters were developed by ECL (Amersham-Pharmacia Biotech).

The results are shown in Figure 5.

Legend of Figure 5:

* indicates prominent tyrosine phosphorylated proteins in stimulated cells.

The results show that incubation of Fc α RI transfectants with anti-Fc α RI Fab A77 induced appearance of several tyrosine-phosphorylated proteins as compared to irrelevant control Fab. The pattern of phosphoproteins appeared identical to the one obtained after multimeric aggregation of Fc α RI, yet differed in its intensity. Similar data were obtained with the Fc α RI_{R209L}/ γ chimeric receptor, while mutations in ITAM abrogated the capacity of this receptor to initiate tyrosine phosphorylation after both monomeric and multimeric targeting.

B) Measurement of cytosolic calcium.

Modulation of the activatory steps was examined regarding effect of anti-Fc α RI Fab A77 on the cytosolic calcium influx ([Ca²⁺]_i), which is a key messenger for cell activation.

1) Experimental protocol

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Aliquots $(1.5 \times 10^6 \text{ cells})$ of human Fc α RI transfectants, or of untransfected RBL-2H3 cells were sensitized with different test reagents (indicated hereafter for each experiment), in complete DMEM supplemented with 20 mM HEPES pH 7.6 during 1 h at 37°C. Cells were then loaded with 4 μ M of the fluorescent probe FURA-2-AM (Molecular probes, Leiden, The Netherlands) for 30 min at 37°C.

After washing, cells were resuspended at 1×10^6 cells/ml in PBS containing 2 mM CaCl₂, 1 mM MgCl₂, 0.5 mg/ml gelatin and placed into a stirred and thermostated bowl. Cells were activated by the addition of 0.1 μ g/ml DNP-HAS antigen (Ag) or 50 nM thapsigargin (Sigma). [Ca²⁺]_i was calculated using the computer software supplied with the spectrofluorimeter (Hitachi F 2000, Salem, New Hampshire) according to the formula

given by GRYNKIEWICZ et al. (J. Biol. Chem. 260: 3440-3450, 1985). No significant cellular auto-fluorescence was observed, and the compounds used did not alter FURA-2-AM fluorescence. The contribution of intracellular stores was determined after stimulation in the presence of 3.5 mM EGTA. For fluorescence quenching studies, $\rm Mn^{2+}$ (200 $\rm \mu M$) was added to cells incubated in $\rm Ca^{2+}$ -free medium (BERTHON and al., Biochem. Pharmacol. 47: 789-794, 1994).

2) Inhibition of the Ca^{2+} plateau phase by anti-Fc α RI Fab

Human Fc α RI transfectants (a, clone 15.4) and untransfected RBL-2H3 cells (b) were sensitized with IgE alone or with IgE plus 10 μ g/ml anti-Fc α RI Fab A77 or irrelevant Fab 320.

The results are shown in Figures 6a and 6b.

Legend of Figures 6a and 6b:

 $a = Human Fc\alpha RI transfectants$

b = untransfected RBL-2H3 cells

Ag: stimulation with DNP-HAS antigen

 \square = IgE alone

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 $O = IgE + anti-Fc\alpha RI Fab A77$

 Δ = IgE + irrelevant Fab 320

Data are representative of at least three separate experiments.

The intracellular calcium peak after FcERI stimulation was unaffected (Fig.6a, \Box), but the plateau phase of elevated [Ca²+]_i was markedly inhibited after preincubation of Fc α RI transfectants with anti-Fc α RI Fab A77 (Fig. 6a, O), as compared to an irrelevant control Fab (Fig. 6a, Δ) or non transfected cells (Fig. 6b).

30 3) Effect of anti-Fc α RI Fab on release from intracellular Ca²⁺ stores

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE alone or with IgE plus 10 μ g/ml anti-Fc α RI Fab A77 or irrelevant Fab 320. Cells were then loaded with FURA-2-AM as indicated and extracellular

calcium was chelated with 3.5 mM EGTA shortly before determining $[{\rm Ca}^{2+}]_i$ to discriminate between calcium release from intracellular stores and calcium entry from the external medium. The results are shown in Figure 6c.

Legend of Figure 6c:

Ag: stimulation with DNP-HAS antigen

 \square = IgE alone

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 $O = IgE + anti-Fc\alpha RI Fab A77$

 Δ = IgE + irrelevant Fab 320

Data are representative of at least three separate experiments.

The results show that anti-Fc α RI Fab A77 treatment had no effect on EGTA-treated cells indicating that it did not inhibit the release of intracellular calcium stores.

4) Anti-FcαRI Fab inhibits Ca2+ influx

To confirm that only calcium influx was affected, external Ca^{2+} was replaced with Mn^{2+} , that enters cells through store-operated calcium channels (SOC) and competes with free internal calcium, thereby quenching FURA-2-AM fluorescence (BERTHON and al., 1994, aforementioned).

Human Fc α RI transfectants (clone 15.4) were sensitized with IgE alone (d) or with IgE plus 10 μ g/ml anti-Fc α RI Fab A77 (e) or irrelevant Fab 320 (f).

The results are shown in Figures 6d-f.

Legend of Figures 6d-f:

Ag: stimulation with DNP-HAS antigen

Mn: addition of Mn²⁺

d = IgE alone

 $e = IgE + anti-Fc\alpha RI Fab A77$

f = IgE + irrelevant Fab 320

Data are representative of at least three separate experiments. The results show that addition of ${\rm Mn}^{2+}$ decreased fluorescence, owing spontaneous entry of ${\rm Mn}^{2+}$ ions into cells. Fc ϵ RI stimulation induced a further

significant decrease in fluorescence as a consequence of Mn^{2+} influx through opened SOC (Fig. 6d). Cell incubation with anti-Fc α RI A77 prior to IgE-dependent stimulation abrogated this effect (Fig. 6e), while an irrelevant Fab 320 was ineffective (Fig. 6f).

Ag induced FURA-2-AM fluorescence quenching, due to $\mathrm{Mn^{2+}}$ influx, with slopes corresponding to calcium entry before and after stimulation are respectively of 2.5 and 4.1 (d), 1.8 and 1.9 (e), and 3.5 (f).

10 5) Anti-FcαRI Fab inhibits events between calcium release from internal stores and the opening of SOC

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To investigate whether FcaRI-mediated inhibition targeted events between calcium release from internal stores and calcium influx, thapsigargin was used, a pharmacologic agent that depletes inositol triphosphate-sensitive stores, resulting in SOC opening, in the absence of transmembrane receptor engagement (THASTRUP and al., Agents Actions 27: 17-23, 1989).

Human Fc α RI transfectants (clone 15.4)(g) and untransfected RBL-2H3 cells (h) were sensitized with 10 μ g/ml anti-Fc α RI Fab A77 or irrelevant Fab 320. After loading the cells with FURA-2-AM, [Ca²⁺]_i was measured following stimulation with 50 nM thapsigargin (Tg).

Data are representative of at least three separate experiments.

The results are shown in Figures 6g and 6h.

Legend of Figures 6g and 6h:

 $g = Human Fc\alpha RI transfectants$

h = untransfected RBL-2H3 cells

Tg: stimulation with thapsigargin

 $O = anti-Fc\alpha RI Fab A77$

 Δ = irrelevant Fab 320

The results show that thapsigargin-induced sustained [Ca $^{2+}$]i elevation was markedly reduced by preincubation with anti-Fc α RI Fab A77 in Fc α RI-

transfected cells , as compared to irrelevant Fab 320 or untransfected cells (Fig. 6h).

EXAMPLE 6: FcαRI TARGETING PREVENTS IGE-MEDIATED MANIFESTATIONS OF ASTHMA *IN VIVO*

The inhibitory activity of Fc α RI being demonstrated in vitro, in vivo targeting of this receptor was tested to know whether it could inhibit inflammatory responses.

not express FcαRI homologs mice do (KABAGAWA and al., Proc. Nat. Acad. Sci. 94: 5261-5266, 10 1997; HAYAMI and al., J. Biol. Chem. 272: 7320-7327, 1997), Balb/c transgenic mice (Tg) expressing the human FcaRI (CD89, line 83) under the control of the CD11b promoter were used, yielding myeloid cell expression similar to that observed in humans (LAUNAY and al., J. 15 Exp. Med. 191: 1999-2009, 2000). Genotyping was done by PCR (LAUNAY and al., 2000, aforementioned). Mice were bred and maintained at the mouse facilities of IFR 02 and Bichat Medical School. All experiments were done in 20 accordance with national guidelines.

Anti-FcaRI Fab immunotherapy was tested in an IgE-mediated animal model of asthma according to ZUBERI and al. (J. Immunol. 164: 2667-2673, 2000) which protocol was adapted. Briefly, FcαRI-transgenic Balb/c mice (Tg) 25 littermate controls (Lt) were immunized and intraperitoneally twice with $10 \mu g$ TNP-OVA (Sigma) 2 mg of aluminium hydroxide gel per 25 g body weight on days 0 and 7. Starting on day 14, mice were challenged intranasally daily for 7 consecutive days with PBS or 2 μg TNP-OVA complexed with 20 μg anti-DNP IgE (IC) 30 the presence of 5 μ g anti-Fc α RI Fab A77 or irrelevant Fab 320. On day 14, mice received 50 μg anti-Fc α RI Fab A77 or control Fab intraperitoneally. Twelve hours after the final intranasal challenge, unrestrained conscious mice were placed in a whole-body plethysmograph chamber (BUXCO 35 Electronics, Sharon, CT). After stabilization for a few

minutes, an aerosol of 300 mM methacholine was delivered for 60 sec.

Changes in airway resistance was calculated every minute for 20 min after methacholine exposure, as follows: enhanced pause (Penh) = [(expiratory time/relaxation time)-1] x (peak expiratory flow/peak inspiratory flow) (ZUANY-AMORIM and al., Science 280: 1265-1267, 1998).

The results are shown in Figure 7a.

10 Legend of Figure 7a:

♦ = Tg PBS

 \bullet = Tg IC + anti-Fc α RI Fab A77

 \triangle = Tg IC + irrelevant Fab 320

 \blacksquare = Lt IC + anti-Fc α RI Fab A77

15 Curves represent mean aiway resistance.

Cumulative areas under the curve (AUC), of corresponding Penh values were means ±SD of three distinct experiments involving at least eight mice per group, and were represented graphically.

The results are shown in Figure.7b.

Legend of Figure 7b:

Tq PBS

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■ = Tg IC + irrelevant Fab 320

 \blacksquare = Tq IC + anti-Fc α RI Fab A77

 \square = Lt IC + anti-Fc α RI Fab A77

* P < 0.05, Student's unpaired t test.

show that results after The repeated intranasal challenge with IgE immune complexes in the presence of an irrelevant Fab, $(Fc\alpha RI^{+})$ Tg mice developed bronchial hyperactivity to inhaled methacholine, as compared to PBS-challenged counterparts (Fig. 7a, 7b). This phenomenon was abrogated by treating transgenic mice (Fig. 7a, 7b). anti-FcαRI Bronchial with Fab hyperactivity was not reduced by anti-FcaRI Fab in (FcαRI) Littermates control non-transgenic (Fig. 7a, 7b).

lung tissue morphological analysis of A sections from FcaRI-transgenic mice was done. Animals were anaesthetized; lungs were inflated by tracheal injection of 1 ml of Optimum Cutter temperature Compound Kingdom), fixed United 4 % Poole, (BDH, paraformaldehyde, dehydrated in graded alcohols, and histopathologic paraffin. Comparative embedded in evaluation of the degree of inflammation was performed on entire H&E-stained lung sections.

The results are shown in Figure 7c-k 10 Legend of Figures 7c-k:

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c-e = control PBS-challenged mice

f-h = antigen-challenged mice treated with irrelevant Fab 320

i-k = antigen-challenged mice treated with 15 anti-Fc α RI Fab A77

Magnification x10 (c,f,i), x100 (d,e,g,h,j,k) Pulmonary histology of antigen-challenged Tg Fab 320 with showed mice treated the irrelevant 20 peribronchial (Fig. 7f) and epithelial (Fig. consisting inflammatory infiltrates mainly granulocytes and mononuclear cells, and diffuse alveolar capillary congestion (Fig. 7h) (see arrows). features were absent in lungs from PBS-challenged mice (Fig. 7c-e) showing normal physiology. Antigen-challenged anti-Fc α RI Fab A77-treated mice showed substantially less inflammation and congestion (Fig. 7i-k). Anti-Fc α RI Fab antigen-induced administration prevented congestion and infiltration by inflammatory cells.

No effects were observed in the lungs of 30 littermates treated with anti-FcaRI Fab (not $(Fc\alpha RI^{-})$ shown).

EXAMPLE 7: EFFECTS OF FCORI TARGETING ON NON-IMMUNE RENAL INFLAMMATION

Anti-FcaRI immunotherapy was also tested after 35 unilateral ureteral obstruction (UUO) in mice,

inflammatory model of interstitial renal fibrosis obstructive nephropathy (KLAHR and MORRISSEY, Am. J. Physiol. Renal Physiol. 283(5): F861-875, 2002). The characterized by tubular dilatation, kidneys are infiltration of inflammatory cells such as macrophages, and epithelial-mesenchymal transition of the kidney. Briefly, unilateral obstruction or the ureter of the left kidney was performed on anaesthetized Tg CD89 mice by ligation at two locations. One day before and daily after chirurgical intervention, mice were treated with either PBS, 100 μ g irrelevant Fab 320 or 100 μ g Fab A77. On day 6, mice were sacrificed and obstructed kidney processed for hispathologic evaluation (Periodic Acid Schiff (PAS) staining and immunohistochemical staining with anti-CD11b antibody).

The results are shown in Figure 8.

Legend of Figure 8:

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Tg CD89 PBS = obstructed kidneys of Tg CD89 mice treated with PBS

Tg CD89 + Fab 320 = obstructed kidneys of Tg CD89 mice treated with irrelevant Fab 320

Tg CD89 + Fab A77 = obstructed kidneys of Tg CD89 mice treated with Fab A77

show PBS-treated kidneys typical The pathologic features of UUO with dilated tubules and 25 cellular infiltration (PAS staining, not shown), notably staining). macrophages (anti-CD11b These typical pathologic features were almost absent in Fab A77-treated cellular infiltration is considerably and decreased. No effects were observed in the kidneys of the 30 Fab 320-treated Tg CD89 mice.